## 510(k) Summary K090742 S00 I

This summary of 510(k) safety and effectiveness information is being submitted in accordance with requirements of 21 CFR Part 807.92.

Date: Jan. 27, 2009

1. Company and Correspondent making the submission:

Name - Samsung Mobile Display Co., Ltd.

Address - San #24, Nongseo-Dong, Giheung-Gu, Yongin-Si, Gyeonggi-Do, Korea, 446-711

Telephone - +82-31-209-3468

Fax - +82-31-209-7369

Contact - Mr. Jooha Park / Senior Manager

Internet - http://www.SAMSUNG.com

2. Device:

Trade/proprietary name

: LTX240AA01-A

**Common Name** 

: Digital Flat Panel X-Ray Detector

Classification Name

: Solid State X-ray Imaging Device

3. Predicate Device:

Manufacturer

: Canon Inc.

**Device** 

: CXDI-50G

510(k) Number

: K031447 (Decision Date - Mar. 26, 2003)

4. Classifications Names & Citations:

21CFR 892.1650, MQB, Solid State X-ray Imaging Device, Class2

- 5. Description:
  - 5.1 General

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Living at the heart of a Digitally enabled society evolving to endless range of applications from Consumer Mobile Display to Medical X-Ray and more to Industrial, Security all beyond our imagination. It's the TFT-based Flat Panel X-Ray Detector that keeps this Digitally based World going forward by providing the most important solution converting transmitted X-Ray into Digital Information. Under the Vision of "Deliver Digitally enriched Images for Visually enabled World!". Now Samsung Mobile Display is thrilled in bringing the first TFT-based Flat Panel X-Ray Detector to this prominent Medical industry.

LTX240AA01-A is a medical image processing unit. Especially, advanced digital imaging process allows considerably efficient diagnosis, all kind of information management, real-time sharing of image information on network.

## 5.2 Product features

LTX240AA01-A is an X-Ray image acquisition device that is based on flat-panel. This device should be integrated with an operating PC and a X-Ray generator. It can do to utilize as digitalizing x-ray images and transfer for radiography diagnostic

#### 6. Indication for use :

LTX240AA01-A Digital Flat Panel X-Ray Detector is indicated for digital imaging solution designed for general radiographic system for human anatomy. It is intended to replace film or screen based radiographic systems in all general purpose diagnostic procedures. Not to be used for mammography.

#### 7. Comparison with predicate device:

Samsung Mobile Display Co., Ltd., believes that the LTX240AA01-A is substantially equivalent to the CXDI-50G of Canon Inc..

## 8. Safety, EMC and Performance Data:

Electrical, mechanical, environmental safety and performance testing according to standard EN/IEC 60601-1 was performed, and EMC testing was conducted in accordance with standard EN/IEC 60601-1-2(2001). All test results were satisfactory.

## 9. Conclusions:

In accordance with the Federal Food, Drug and Cosmetic Act, 21 CFR Part 807 and based on the information provided in this premarket notification Samsung Mobile Display Co., Ltd. concludes that The LTX240AA01-A is safe and effective and substantially equivalent to predicate devices as described herein.

10. Samsung Mobile Display Co., Ltd. will update and include in this summary any other information deemed seasonably necessary by the FDA.

**END** 



Public Health Service

Food and Drug Administration 10903 New Hampshire Avenue Document Control Room – WO66-G609 Silver Spring, MD 20993-0002

Samsung Mobile Display Co., Ltd.

% Mr. Vince Lee

Manager

E-WOO Technology USA, Inc.

256 North Sam Houston Pkwy E., Suite 115

**HOUSTON TX 77060** 

AUG 2 3 2013

Re: K090742

Trade/Device Name: Digital Flat Panel X-Ray Detector/LTX240AA01-A

Regulation Number: 21 CFR 892.1680 Regulation Name: Stationary x-ray system

Regulatory Class: II Product Code: MQB Dated: September 1, 2009 Received: September 11, 2009

## Dear Mr. Lee:

This letter corrects our substantially equivalent letter of September 18, 2009.

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

If your device is classified (see above) into class II (Special Controls), it may be subject to such additional controls. Existing major regulations affecting your device can be found in Title 21, Code of Federal Regulations (CFR), Parts 800 to 895. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Parts 801 and 809); medical device reporting (reporting of

medical device-related adverse events) (21 CFR 803); and good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820). This letter will allow you to begin marketing your device as described in your Section 510(k) premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus, permits your device to proceed to the market.

If you desire specific advice for your device on our labeling regulation (21 CFR Parts 801 and 809), please contact the Office of *In Vitro* Diagnostic Device Evaluation and Safety at (301) 796-5450. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <a href="http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm">http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm</a> for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address <a href="http://www.fda.gov/cdrh/industry/support/index.html">http://www.fda.gov/cdrh/industry/support/index.html</a>.

Sincercity 10ms,

Janine M. Morri

Acting Director

Division of Radiological Devices
Office of In Vitro Diagnostic Device

**Evaluation and Safety** 

Center for Devices and Radiological Health

**Enclosure** 

# Indications for Use

		510(k) Number(if known): K	09074	1000	
		Device Name: Digital Flat Panel X-Ray Detector /LTX240AA01-A			
		Indications for Use:			
	•	LTX240AA01-A Digital F imaging solution design anatomy. It is intended to in all general purpose mammography.	replace film or	al radiographic sy screen based radi	stem for human ographic systems
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		Prescription Use	: ************************************		
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